## Generalization in Deep Learning

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#### Abstract

This paper explains why deep learning can generalize well, despite large capacity and possible algorithmic instability, nonrobustness, and sharp minima, effectively addressing an open problem in the literature. Based on our theoretical insight, this paper also proposes a family of new regularization methods. Its simplest member was empirically shown to improve base models and achieve state-of-theart performance on MNIST and CIFAR-10 benchmarks. Moreover, this paper presents both data-dependent and data-independent generalization guarantees with improved convergence rates. Our results suggest several new open areas of research.

## 1 Introduction

Deep learning has been a great practical success and had a profound impact on the machine learning and artificial intelligence literature. Along with its practical success, theoretical properties of deep learning have been investigated in the literature. For expressivity of neural networks, there are classical results of universality (Leshno et al., 1993) and the exponential advantage against hand-crafted features (Barron, 1993). Another series of theoretical studies have considered how trainable (or optimizable) deep model classes are, revealing their nontrivial properties beyond non-convexity (Choromanska et al., 2015; Safran and Shamir, 2016; Kawaguchi, 2016; Yun et al., 2017). On the other hand, having an expressive and trainable model class does not by itself imply a good performance in predicting the future, owing to possible over-fitting to training data. This leads to the study of *generalization*, which is the focus of this paper.

Some classical theory work attributes generalization

ability to the use of small-capacity model classes (Mohri et al., 2012). From the viewpoint of compact representation related to small capacity, it has been shown that deep model classes have an exponential advantage to represent certain natural target functions when compared to shallow model classes (Pascanu et al., 2014; Montufar et al., 2014; Livni et al., 2014; Telgarsky, 2016; Poggio et al., 2017). In other words, when some assumptions implicit in the model class (e.g. deep composition of piecewise linear transformations) are approximately satisfied by the target function, one can get very good generalization, compared to methods which do not rely on that assumption. However, a recent paper (Zhang et al., 2017a) has empirically shown that successful deep model classes have sufficient capacity to memorize random labels. This observation has been called an "apparent paradox" and led to active discussions by many researchers (Arpit et al., 2017; Zhang et al., 2017b; Krueger et al., 2017; Hoffer et al., 2017; Wu et al., 2017; Neyshabur et al., 2017; Dziugaite and Roy, 2017; Dinh et al., 2017). Dinh et al. (2017) conclude that explaining why deep learning models can generalize well despite their overwhelming capacity is an open area of research.

In this paper, we propose an explanation for this "apparent paradox" to address the open problem of explaining why deep learning can sometimes generalize well despite its large capacity and possible instability, nonrobustness, and sharp minima. In Section 3, we theoretically show the existence of the same "apparent paradox" not only for deep learning but also for machine learning in general. Our resolution of the apparent paradox sheds light on the need to rethink generalization and learning theory. As a result of such a rethinking, we obtain an explanation for generalization in deep learning in Section 4, an improvement of generalization bounds in Section 5, and a useful theoretical insight for regularization in Section 6.

## 2 Preliminaries

In this section, we define the notation and briefly review the relevant generalization theory based on model complexity (e.g., Rademacher complexity and VC di-

mension), stability, robustness, and flat minima.

#### 2.1 Notation

Let  $x \in \mathcal{X}$  be an input and  $y \in \mathcal{Y}$  be a target. Let m be the size of a training dataset. Let  $f_{\mathcal{A}(S_m)} : \mathcal{X} \to \mathcal{Y}$  be a model learned by a learning algorithm  $\mathcal{A}$  (including random seeds for simplicity) with a training dataset  $S_m = \{(x_1, y_1), \ldots, (x_m, y_m)\}$ . We assume that  $S_m$  is generated by i.i.d. draws unless otherwise specified.

Let  $\ell$  be a loss function. Let R[f] and  $\hat{R}_m[f]$  be the expected risk and empirical risk of a function f as  $R[f] = \mathbb{E}_{x,y \sim P_{(x,y)}}[\ell(f(x),y)]$  and  $\hat{R}_m[f] = \frac{1}{m} \sum_{i=1}^m \ell(f(x_i),y_i)$ , where  $(x_i,y_i) \sim P_{(x,y)}$ . Let  $\hat{R}_i[f] = \ell(f(x_i),y_i)$ . We define  $\hat{R}_{\text{val}}$  and  $\hat{R}_{\text{val},i}$  in the same way for a held-out validation dataset. Let F be a set of functions, a model class, or a hypothesis set. Let  $\mathcal{L}_F$  be a family of loss functions associated with F, defined by  $\mathcal{L}_F = \{g: f \in F, g(x,y) \triangleq \ell(f(x),y)\}$ . Let  $w \in \mathbb{R}^n$  be the trainable parameters for a parameterized model class. All vectors are column vectors in this paper.

Let  $h^{(l)}(x)$  be the pre-activation vector of the l-th hidden layer. Let H be the number of hidden layers. For example,  $f_{\text{out}}(x) = \operatorname{softmax}(h^{(H+1)}(x))$  and  $f(x) = \operatorname{argmax}(h^{(H+1)}(x))$  are a typical model output and prediction for classification. With such  $f(x) = \operatorname{argmax}(h^{(H+1)}(x))$  for classification, let  $\hat{R}_{m,\rho}[f]$  be the usual empirical margin loss of f (see Appendix A for detail). Let  $\lambda_{\max}(M)$  and  $\|M\|_2$  be the largest singular value and the spectral norm of a matrix M.

## 2.2 Overview of Generalization Theory

A goal in machine learning is typically framed as the minimization of the expected risk  $R[f_{\mathcal{A}(S_m)}]$ . We usually aim to minimize the non-computable expected risk  $R[f_{\mathcal{A}(S_m)}]$  by minimizing the computable empirical risk  $\hat{R}_m[f_{\mathcal{A}(S_m)}]$  (i.e., empirical risk minimization). A goal of generalization theory is to explain and justify why and how minimizing  $\hat{R}_m[f_{\mathcal{A}(S_m)}]$  is a sensible approach to minimize  $R[f_{\mathcal{A}(S_m)}]$  by analyzing

the generalization gap 
$$\triangleq R[f_{\mathcal{A}(S_m)}] - \hat{R}_m[f_{\mathcal{A}(S_m)}].$$

A challenge of analyzing the generalization gap stems from the dependence of  $f_{\mathcal{A}(S_m)}$  on the same dataset  $S_m$  used in the definition of  $\hat{R}_m$ . Several approaches in (statistical) generalization theory have been developed to handle this dependence in various ways.

The model complexity approach handles this dependence by decoupling  $f_{\mathcal{A}(S_m)}$  from the particular  $S_m$  as

$$R[f_{\mathcal{A}(S_m)}] - \hat{R}_m[f_{\mathcal{A}(S_m)}] \le \sup_{f \in F} R[f] - \hat{R}_m[f],$$

and by carefully analyzing the right-hand side. Because the cardinality of F is typically (uncountably) infinite,

a direct use of the union bound over all elements in F gives a vacuous bound, leading to the need to consider different quantities to characterize F; e.g., model complexities of F, such as Rademacher complexity and VC dimension. For example, if the codomain of  $\ell$  is in [0,1], we have (Mohri et al., 2012, Theorem 3.1) that for any  $\delta > 0$ , with probability at least  $1 - \delta$ ,

$$\sup_{f \in F} R[f] - \hat{R}_m[f] \le 2\Re_m(\mathcal{L}_F) + \sqrt{\frac{\ln \frac{1}{\delta}}{2m}},$$

where  $\mathfrak{R}_m(\mathcal{L}_F)$  is the Rademacher complexity of  $\mathcal{L}_F$ , which then can be bounded by the Rademacher complexity of F,  $\mathfrak{R}_m(F)$ . However, for the deep learning model classes F, the known bounds on  $\mathfrak{R}_m(F)$  explicitly grow exponentially in depth (Sun et al., 2016; Neyshabur et al., 2015; Xie et al., 2015). Another type of bound explicitly grows linearly in the number of trainable parameters (Shalev-Shwartz and Ben-David, 2014). Both types of bounds cannot account for the practical observations with deep learning.

The stability approach deals with the dependence of  $f_{\mathcal{A}(S_m)}$  on the dataset  $S_m$  by considering the stability of algorithm  $\mathcal{A}$  with respect to datasets. The considered stability is a measure of how much changing a data point in  $S_m$  can change  $f_{\mathcal{A}(S_m)}$ . For example, if the algorithm  $\mathcal{A}$  has uniform stability  $\beta$  (w.r.t.  $\ell$ ) and if the codomain of  $\ell$  is in [0, M], we have (Bousquet and Elisseeff, 2002) that for any  $\delta > 0$ , with probability at least  $1 - \delta$ ,

$$R[f_{\mathcal{A}(S_m)}] - \hat{R}_m[f_{\mathcal{A}(S_m)}] \le 2\beta + (4m\beta + M)\sqrt{\frac{\ln\frac{1}{\delta}}{2m}}.$$

Based on previous work on stability (e.g., Hardt et al. 2015; Kuzborskij and Lampert 2017; Gonen and Shalev-Shwartz 2017), one may conjecture the reason of generalization in deep learning, the proving of which is still an open problem.

The robustness approach successfully avoids dealing with some details of the dependence of  $f_{\mathcal{A}(S_m)}$  on  $S_m$  by considering the robustness of algorithm  $\mathcal{A}$  for all possible datasets. In contrast to stability, which measures the variation w.r.t. data samples, robustness is the measure of how much the loss value can vary w.r.t. the input space of (x,y). As a result, this approach can naturally take the dependence on a particular  $S_m$  into consideration to some degree. For example, if algorithm  $\mathcal{A}$  is  $(\Omega, \epsilon(\cdot))$ -robust and the codomain of  $\ell$  is upper-bounded by M, given a dataset  $S_m$ , we have (Xu and Mannor, 2012) that for any  $\delta > 0$ , with probability at least  $1 - \delta$ ,

$$|R[f_{\mathcal{A}(S_m)}] - \hat{R}_m[f_{\mathcal{A}(S_m)}]| \le \epsilon(S_m) + M\sqrt{\frac{2\Omega \ln 2 + 2\ln \frac{1}{\delta}}{m}}$$

The robustness approach requires an a priori known and fixed partition of the input space such that the

number of sets in the partition is  $\Omega$  and the change of loss values in each set in the partition is bounded by  $\epsilon(S_m)$  for all  $S_m$  (Definition 2 and the proof of Theorem 1 in Xu and Mannor 2012). In classification, if the margin is ensured to be large, we can fix the partition with balls of radius corresponding to the large margin, filling the input space. Recently, this idea was applied for deep learning (Sokolic et al., 2017a,b). These previous results still suffer from the curse of (a priori known effective) dimensionality of the input space, which is an important open problem.

Flat minima also correspond to a large margin as well as robustness in parameter space (as opposed to robustness in the input space). Several studies have provided arguments for generalization in deep learning based on flat minima (Keskar et al., 2017; Zhang et al., 2017b). However, Dinh et al. (2017) showed that flat minima in practical deep learning model classes can be turned into sharp minima via re-parameterization without changing the generalization gap, and hence it requires further investigation.

# 3 Rethinking Generalization in Machine Learning

Zhang et al. (2017a) empirically showed that several deep model classes can memorize random labels, while having the ability to produce zero training error and small test errors for particular natural datasets (e.g., CIFAR-10). They also empirically observed that the regularization on the norm of weights seemed to be unnecessary to obtain small test errors, in contradiction to "traditional wisdom."

Supporting and extending these *empirical* observations, our Theorem 8 in Appendix B.1 theoretically proves that the class of over-parameterized linear models can memorize any training data and decrease the training and test errors arbitrarily close to zero (including zero) with the norm of parameters being arbitrarily large, even when the parameters are arbitrarily far from the ground-truth parameters. Furthermore, Corollary 9 in Appendix B.2 shows that traditional wisdom on the norm of the parameter w can fail to explain generalization even in linear models. All proofs in this paper are presented in the appendix.

Proposition 1 states that these phenomena are not limited to deep learning and linear model classes: any machine learning model class essentially has the core property of these phenomena. Proposition 1 can be rephrased more essentially as follows. The actual value of  $R[f] - \hat{R}[f]$  is clearly determined by what we get as f, independent of the process used to get f (learning mechanism  $\mathcal{A}$ ) and from which set we choose f (model class F and flat-minima). In contrast, the traditional wisdom based on the current generalization

theory states that how we get f (learning mechanism A) and the set from which we choose f (model class F or flat-minima) are all that matters. This has created the "apparent paradox" in the literature.

**Proposition 1** Given an (unknown) measure  $P_{(x,y)}$  and a dataset  $S_m$ , suppose that there exists  $f_{\epsilon}^*$  for some  $\epsilon \geq 0$  such that  $R[f_{\epsilon}^*] - \hat{R}_m[f_{\epsilon}^*] \leq \epsilon$ . Then,

- (i) For any model class F whose model complexity is large enough to memorize any dataset and which includes  $f_{\epsilon}^*$  possibly at an arbitrarily sharp minimum, there exists  $(A, S_m)$  such that the generalization gap is at most  $\epsilon$ , and
- (ii) For any dataset  $S_m$ , there exist arbitrarily unstable and arbitrarily non-robust algorithms A such that the generalization gap of  $f_{A(S_m)}$  is at most  $\epsilon$ .

#### 3.1 Demystifying the Generalization Puzzle

A recent paper (Dinh et al., 2017) states that flat minima can be turned into sharp minima, resulting in a "contradiction" of the flat minima approach in generalization theory. From Proposition 1, it is now clear that the apparent "contradiction" is not limited to sharp minima in deep learning, but is rather more fundamental. This section carefully looks at the meaning of mathematical statements, in order to clarify what has been considered to be paradoxical results in the literature (e.g., Arpit et al. 2017; Zhang et al. 2017b; Dinh et al. 2017).

Generalization theory can be considered to provide two types of statements relevant to the scope of this paper. The first type (which comes from upper bounds) is logically in the form of "p implies q," where p:= "the model complexity is small" (or another statement about stability, robustness, or flat minima), and q:= "the generalization gap is small." Notice that "p implies q" does not imply "q implies p." Thus, based on statements of this type, it is entirely possible that the generalization gap is small even when the model complexity is large or the learning mechanism is unstable, non-robust, or subject to sharp minima.

The second type (which comes from lower bounds) is logically in the following form: in a set  $U_{\rm all}$  of all possible problem configurations, there exists a subset  $U\subseteq U_{\rm all}$  such that "q implies p" in U (with the same definitions of p and q as in the previous paragraph). For example, Mohri et al. (2012, Section 3.4) derived lower bounds on the generalization gap by showing the existence of a "bad" distribution that characterizes U. However, if the problem instance at hand (e.g., object classification with MNIST or CIFAR-10) is not in such a U in the proofs (e.g., if the data distribution is not among the "bad" ones considered in the proofs), q does not necessarily imply p. Thus, it is still naturally possible that the generalization gap is small with large model

complexity, instability, non-robustness, and sharp minima. Therefore, there is no contradiction or paradox.

#### 3.2 Practical Role of Generalization Theory

From the previous two sections, we can see that there is a logically expected difference between the scope in theory and the focus in practice; it is logically expected that there are problem instances where theory does not make claims about generalization observable in practice. In order for generalization theory to have maximal impact, we must be clear on a set of different roles it can play, and then work to extend and strengthen it in each of these roles. We have identified the following practical roles for theory (which are further explained in Appendix B.4):

Role 1 Provide guarantees on expected risk.

Role 2 Guarantee generalization gap

Role 2.1 to be small for a given fixed  $S_m$ , and/or Role 2.2 to approach zero with a fixed model class as m increases.

Role 3 Provide theoretical insights to guide the search over model classes.

Previous attempts to explain generalization in deep learning can be seen as trying to achieve all Roles 1-3 of the theory at the same time, which might be too ambitious, considering the expected logical difference between theory and practice. For example, model complexity might be a good theoretical insight to guide the model class search, but it would not be tight enough to explain generalization itself. The rest of this paper presents generalization theory that separately addresses each role: Role 1 (Section 4), Role 2 (Section 5) and Role 3 (Section 6).

## 4 Understanding Generalization in Practical Paradigm

Role 1 of the theory is to provide guarantees on a quantity that we cannot compute: the expected risk. Although it is often ignored, we can compute the validation error as well as the training error in practice. That is, in practical deep learning, we typically adapt the training-validation paradigm, usually with a held-out validation set. We then search over the model classes by changing architectures (and other hyper-parameters) to obtain a low validation error. In this view, the reason why deep learning can generalize well becomes clear: we can generalize well because we can obtain a good model via model search with validation errors. Indeed, as an example, Proposition 2 states that if a validation error is small (which is usually ensured in practice), it is guaranteed to generalize well, independently of the capacity, stability, robustness, and flat minima. An achievement of a small validation error will typically happen when the target function is consistent with the assumptions implicit in the training objective and model class. We can thus end up with a very different effective capacity (as selected by model search using the validation set) depending on whether the training data are random or have interesting structure which the neural network can capture.

**Proposition 2** (example of generalization guarantee via validation error) Let  $z_{f,i} = R[f] - \hat{R}_{\text{val},i}[f]$ . Suppose that  $\mathbb{E}[z_{f,i}^2] \leq \gamma^2$  and  $|z_{f,i}| \leq C$  almost surely, for all  $(f,i) \in F_{\text{val}} \times \{1,\ldots,m_{\text{val}}\}$ . Then, for any  $\delta > 0$ , with probability at least  $1 - \delta$ , the following holds for all  $f \in F_{\text{val}}$ :

 $R[f] \le \hat{R}_{\text{val}}[f] + \frac{2C \ln(\frac{|F_{\text{val}}|}{\delta})}{3m_{\text{val}}} + \sqrt{\frac{2\gamma^2 \ln(\frac{|F_{\text{val}}|}{\delta})}{m_{\text{val}}}}.$ 

Here,  $F_{\rm val}$  is defined to be a finite set of models f that are proposed independently of a held-out validation dataset. To be independent of a validation dataset,  $F_{\rm val}$  must be fixed without the influence of the validation dataset. In other words,  $|F_{\rm val}|$  is the number of times we use the validation dataset in our decision-making to choose a final model. For example,  $F_{\rm val}$  can be a set of models f such that each element f is a result at the end of each epoch during training with at least 99.5% training accuracy. In this example,  $|F_{\rm val}|$  is at most (the number of epochs)  $\times$  (the number of hyperparameter searches), and is likely much smaller than that owing to the 99.5% training accuracy criteria.

The bound in Proposition 2 is non-vacuous and tight enough to be practically meaningful. For example, consider a classification task with 0–1 loss. Set  $m_{\rm val}=10,000$  (e.g., MNIST and CIFAR-10) and set  $\delta=0.1$ . Then, even in the worst case with C=1 and  $\gamma^2=1$  and even with  $|F_{\rm val}|=1,000,000,000$ , we have with probability at least 0.9 that  $R[f] \leq \hat{R}_{\rm val}[f] + 6.94\%$  for all  $f \in F_{\rm val}$ . In a non-worst-case scenario, for example, with C=1 and  $\gamma^2=(0.05)^2$ , we can replace 6.94% by 0.49%. With a larger validation set (e.g., as in ImageNet) and/or more optimistic C and  $\gamma^2$ , we can obtain much better bounds.

Therefore, bounds on expected risk via validation errors such as Proposition 2 can explain why deep learning has the ability to generalize well, despite the large capacity and possible instability, nonrobustness, and sharp minima. Meanwhile, such reasoning with Proposition 2 suggests another open question: why is it that we can often find good architectures (and other hyper-parameters) that get low validation errors (with non-exponentially-many trials relative to the validation dataset size, as in  $\ln |F_{\rm val}|/m_{\rm val}$  in Proposition 2). A close look at the deep learning literature seems to suggest that this question is fundamentally related to the process of science and engineering, because many successful architectures have been designed based on the

physical properties and engineering priors of the problems at hand (e.g., hierarchical nature, convolution, architecture for motion such as that by Finn et al. 2016, memory networks, and so on).

Although Proposition 2 poses a concern of increasing the generalization bound when using a single validation dataset with too large  $F_{\rm val}$ , the rate of increase is only  $\ln |F_{\rm val}|$  and  $\sqrt{\ln |F_{\rm val}|}$ . This rate seems to be slow enough for the current deep learning literature. For example, even a very large number  $e^{100} \approx 2.69 \times 10^{43}$  of the uses of a validation dataset incurs only  $\ln |F_{\rm val}| = 100$  relative to  $m_{\rm val}$ . Nevertheless, the next section presents different generalization bounds that can avoid this possible concern (also see Ng 1997; Cawley and Talbot 2010 for more discussions on the possible over-fitting to validation dataset).

## 5 Generalization Guarantees Specific to Neural Networks

The previous section provided an explanation for generalization in practical situations. However, it is still of great interest to obtain theoretical guarantees of the generalization gap,  $R[f_{\mathcal{A}(S_m)}] - \hat{R}_m[f_{\mathcal{A}(S_m)}]$  (Role 2), which is the focus of this section. To closely analyze neural networks, this section presents a direct analysis of neural networks rather than deriving results about neural networks from more generic theory based on capacity, stability, or robustness.

#### 5.1 Model Description via Deep Paths

We consider general neural networks with any depth that have the structure of a directed acyclic graph (DAG) with ReLU nonlinearity and/or max pooling. (e.g., any feedforward network or roll-out of a recurrent network with convolutional and/or fully connected layers). From the DAG structure, we can write the value of each output unit as the summation over all values of all paths from the input to the k-th output unit; that is, for all  $k \in \{1, \ldots, d_y\}$ ,

$$h_k^{(H+1)}(x) = \sum_{\text{path}} \bar{x}_{\text{path}} \bar{\sigma}_{\text{path}}(x, w_\sigma) \bar{w}_{\text{path},k}$$
$$= [\bar{x} \circ \bar{\sigma}(x, w_\sigma)]^\top \bar{w}_k, \tag{1}$$

where  $\sum_{\text{path}}$  represents the summation over all paths from the input to the k-th output unit. Here,  $[\bar{x} \circ \bar{\sigma}(x, w_{\sigma})]$  and  $\bar{w}_k$  are the vectorized versions of the summation with the vector size being the number of the paths. Additionally,  $\bar{\sigma}(x, w_{\sigma})$  represents the activation of each path, corresponding to ReLU nonlinearity and max pooling;  $\bar{\sigma}_j(x, w_{\sigma}) = 1$  if the j-th path is active, and  $\bar{\sigma}_j(x, w_{\sigma}) = 0$  otherwise. Each entry of the vector  $\bar{w}$  is the product of the weight parameters for each path, where vectors  $w_{\sigma}$  and  $\bar{w}_k$  share the same weight parameters. We use  $[\bar{x} \circ \bar{\sigma}(x, w_{\sigma})]$  to denote the element-wise

product of  $\bar{x}$  and  $\bar{\sigma}(x, w_{\sigma})$ . See Appendix D.2 for an example of equation (1).

## 5.2 Data-dependent Guarantee

We first focus on regression with squared loss, for which we present our direct analysis. Let  $z_i = [\bar{x}_i \circ \bar{\sigma}(x_i, w_{\sigma})]$  (note the dependence on x and  $w_{\sigma}$ ). Let  $\bar{w}_{\mathcal{A}(S_m),k}$  be  $\bar{w}_k$  obtained by algorithm  $\mathcal{A}$  with dataset  $S_m$ . Then, from equation (1), we can decompose the generalization gap into three terms as

$$R[f_{\mathcal{A}(S_m)}] - \hat{R}_m[f_{\mathcal{A}(S_m)}]$$

$$= \sum_{k=1}^{d_y} \left[ \bar{w}_{\mathcal{A}(S_m),k}^{\top} \left( \mathbb{E}[zz^{\top}] - \frac{1}{m} \sum_{i=1}^m z_i z_i^{\top} \right) \bar{w}_{\mathcal{A}(S_m),k} \right]$$

$$+ 2 \sum_{k=1}^{d_y} \left[ \left( \frac{1}{m} \sum_{i=1}^m y_{ik} z_i^{\top} - \mathbb{E}[y_k z^{\top}] \right) \bar{w}_{\mathcal{A}(S_m),k} \right]$$

$$+ \mathbb{E}[y^{\top}y] - \frac{1}{m} \sum_{i=1}^m y_i^{\top}y_i.$$

Each of the three terms contains a difference between a sum of *dependent* random variables and its expectation. Accordingly, we want a dataset  $S_m$  to be "good" in the sense that these differences are small, as defined as follows.

**Definition 3** (concentrated dataset) A dataset  $S_m$  is said to be  $(\beta_1, \beta_2, \beta_3)$ -concentrated with respect to the square loss and  $w_{\sigma}$  if  $\lambda_{\max} \left( \mathbb{E}[zz^{\top}] - \frac{1}{m} \sum_{i=1}^{m} z_i z_i^{\top} \right) \leq \beta_1$ ,  $\left\| \frac{1}{m} \sum_{i=1}^{m} y_{ik} z_i - \mathbb{E}[y_k z] \right\|_{\infty} \leq \beta_2$ , and,  $\mathbb{E}[\|y\|_2^2] - \frac{1}{m} \sum_{i=1}^{m} \|y_i\|_2^2 \leq \beta_3$ .

Here, we can see the benefit of deep learning from the viewpoint of "deep-path" feature learning; even if a given  $S_m$  is not concentrated in the original space, optimizing  $w_{\sigma}$  can result in the concentration in the space of z, defined by the paths of deep networks.

Proposition 4 states that if a given dataset  $S_m$  is good in the sense of Definition 3, then the generalization gap is deterministically bounded based only on the quality of what we get as w, independently of how we choose w (stability and robustness) and the set from which we select w (model complexity and flat minima), as desired (Role 2.1).

**Proposition 4** (data-dependent and deterministic bound) Suppose that  $S_m$  is  $(\beta_1, \beta_2, \beta_3)$ -concentrated with respect to the square loss and  $w_{\sigma}$ . Then, deterministically,

$$R[f_{\mathcal{A}(S_m)}] \leq \hat{R}_m[f_{\mathcal{A}(S_m)}] + \beta_1 \sum_{k=1}^{d_y} \|\bar{w}_{\mathcal{A}(S_m),k}\|_2^2 + \beta_2 \sum_{k=1}^{d_y} \|\bar{w}_{\mathcal{A}(S_m),k}\|_1 + \beta_3.$$

#### 5.3 Data-independent guarantee

A remaining question is whether we can guarantee a randomly drawn dataset to be "good" in the sense of the concentration. Notice that if  $z_i$  were independent of each other, it would have been very easy to guarantee that. However, the  $z_i$  are dependent owing to the dependence of  $w_{\sigma}$  on  $S_m$ . We then observe that each step of the (stochastic) gradient decent greedily chooses the best direction only in terms of  $\bar{w}$ , but not in terms of  $w_{\sigma}$  (see Appendix D.3). This observation leads to a hypothesis that learning a good  $w_{\sigma}$  does not require all of the information contained in  $S_m$ , and hence the dependence of  $z_i$  is not entirely "bad."

## 5.3.1 Empirical Observations

As a first step to investigate the dependence of  $z_i$ , we evaluated the following novel two-phase training procedure that explicitly breaks the dependence. In this procedure, we first train a network in a standard way, but only with partial training dataset of  $\alpha m$  size, where  $\alpha \in (0,1)$  (standard phase). We then freeze  $w_{\sigma}$  and train only  $\bar{w}$  with the whole training dataset of size m (freeze phase). Note that vectors  $w_{\sigma}$  and  $\bar{w}$  contain the same shared parameters in the standard phase, whereas these parameters are untied in freeze phase. See Appendix D.4 for a simple implementation of this two-phase training procedure that requires at most (approximately) twice as much computational cost as the normal training procedure.

We implemented the two-phase training procedure with the MNIST and CIFAR-10 datasets. Because we are not aiming for the state-of-the-art results in this section, we used computationally less expensive settings when compared with the next section (where we achieve better performance). The test accuracies of the standard training procedure (base case) were 99.47% for MNIST (ND), 99.72% for MNIST, and 92.89% for CIFAR-10. MNIST (ND) indicates MNIST with no data augmentation. The experimental details are in Appendix D.5. Our source code is available at:

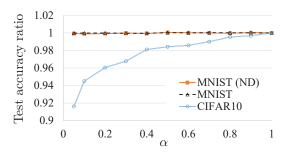


Figure 1: Test accuracy ratio (Two-phase/Base). Notice that the y-axis starts with already high accuracy, even with very small data size  $\alpha m$  for learning  $w_{\sigma}$ .

#### http://lis.csail.mit.edu/code/gdl.html

Figure 1 shows the test accuracy ratio for varying  $\alpha$ : the test accuracy of the standard training procedure divided by the test accuracy of this two-phase training procedure. The plot in Figure 1 starts with  $\alpha=0.05$ , for which  $\alpha m=3000$  in MNIST and  $\alpha m=2500$  in CIFAR-10. Somewhat surprisingly, using a much smaller dataset for learning  $w_{\sigma}$  still resulted in competitive performance. Notably, a dataset on which we can more easily get a better generalization (i.e., MNIST) allowed us to use smaller  $\alpha$  to achieve competitive performance. This is consistent with our discussion above.

## 5.3.2 Theoretical Results

Our two-phase training procedure (standard phase plus freeze phase) forces  $z_{\alpha m+1}, \ldots, z_m$  to be independent random variables, while maintaining competitive practical performance. Thus, by using this two-phase training procedure, one can obtain the following guarantees on practical models, addressing some issues left open in previous studies. Let  $m_{\sigma} = (1 - \alpha)m$  and  $S_{m_{\sigma}} = \{(x_{\alpha m+1}, y_{\alpha m+1}), \ldots, (x_m, y_m)\}.$ 

**Assumption 1** Let  $A^{(i)} = \mathbb{E}_x[zz^{\top}] - z_i z_i^{\top}$ ,  $B_{kk'}^{(i)} = y_{ik} z_{i,k'} - \mathbb{E}_{x,y}[y_k z_{k'}]$ , and  $R^{(i)} = \mathbb{E}_y[||y||_2^2] - ||y_i||_2^2$ . Assume that for all  $i \in \{\alpha m + 1, ..., m\}$ ,

- $C_{zz} \ge \lambda_{\max}(A^{(i)})$  and  $\gamma_{zz}^2 \ge \|\mathbb{E}_x[(A^{(i)})^2]\|_2$
- $C_{yz} \ge \max_{k,k'} |B_{kk'}^{(i)}| \text{ and } \gamma_{yz}^2 \ge \max_{k,k'} \mathbb{E}_x[(B_{kk'}^{(i)})^2])$
- $C_y \ge |R^{(i)}|$  and  $\gamma_y^2 \ge \mathbb{E}_x[(R^{(i)})^2]$ .

**Lemma 5** Suppose that Assumption 1 holds, and use its notation. Set  $\beta_1 = \frac{2C_{zz}}{3m_\sigma} \ln \frac{3d_z}{\delta} + \sqrt{\frac{2\gamma_{zz}^2}{m_\sigma}} \ln \frac{3d_z}{\delta}$ ,  $\beta_2 = \frac{2C_{yz}}{3m_\sigma} \ln \frac{6d_yd_z}{\delta} + \sqrt{\frac{\gamma_{yz}^2}{m_\sigma}} \ln \frac{6d_yd_z}{\delta}$ , and  $\beta_2 = \frac{2C_y}{3m_\sigma} \ln \frac{3}{\delta} + \sqrt{\frac{2\gamma_y^2}{m_\sigma}} \ln \frac{3}{\delta}$ . Then, for any  $\delta > 0$ , with probability at least  $1 - \delta$ , a dataset  $S_{m_\sigma}$  is  $(\beta_1, \beta_2, \beta_3)$ -concentrated with respect to the squared loss and a frozen  $w_\sigma$ .

**Theorem 6** (probabilistic bound via random matrix theory) Suppose that Assumption 1 holds. Set  $(\beta_1, \beta_2, \beta_3)$  to be those in Lemma 5. Consider any algorithm  $\mathcal{A}$  such that if  $S_{m_{\sigma}}$  is  $(\beta_1, \beta_2, \beta_3)$ -concentrated with respect to the squared loss and a frozen  $w_{\sigma}$ ,  $\sum_{k}^{d_y} \|\bar{w}_{\mathcal{A}(S_m),k}\|_1 \leq C_{w1}$  and  $\sum_{k}^{d_y} \|\bar{w}_{\mathcal{A}(S_m),k}\|_2^2 \leq C_{w2}$  (otherwise, the norms need not be bounded). Then, for any  $\delta > 0$ , with probability at least  $1 - \delta$ .

$$R[f_{\mathcal{A}(S_m)}] \le \hat{R}_{m_{\sigma}}[f_{\mathcal{A}(S_m)}] + C_{w2}\beta_1 + C_{w1}\beta_2 + \beta_3.$$

Theorem 6 only requires the norms to be bounded for "good" datasets, allowing unbounded norms for "bad" datasets. For classification with 0–1 loss, we adapt the model complexity approach for the two-phase training procedure, yielding Theorem 7.

**Theorem 7** Fix  $\rho > 0$ . Suppose that  $\mathbb{E}_x[\|\bar{x} \circ \bar{\sigma}(x, w_{\sigma})\|_2^2] \leq C_{\sigma}^2$  and  $\max_k \|\bar{w}_k\|_2 \leq C_w$  for all  $f \in F$ . Then, for any  $\delta > 0$ , with probability at least  $1 - \delta$ , the following holds for all  $f \in F$ :

$$R[f] \le \hat{R}_{m_{\sigma},\rho}[f] + \frac{2d_y^2(1-\alpha)^{-1/2}C_{\sigma}C_w}{\rho\sqrt{m}} + \sqrt{\frac{\ln\frac{1}{\delta}}{2m_{\sigma}}}.$$

Theorems 6 and 7 provide the first generalization bounds for practical deep learning models that do not necessarily have dependence on the number of weights (unlike previous VC dimension approaches), and exponential dependence on depth (unlike Sun et al. 2016; Neyshabur et al. 2015; Xie et al. 2015) and effective input dimensionality (unlike Sokolic et al. 2017a,b). For example, the comparable parts to  $(1-\alpha)^{-1/2}C_{\sigma}C_{w}$  in our Theorems 7 are in the form of  $2^H C_X \prod_{\ell=1}^{H+1} C_{W^{(\ell)}}$ in previously known bounds where  $C_X$  and  $C_{W^{(\ell)}}$  correspond to the upper-bounds on the norms of the input x and of the weight matrix at the  $\ell$ -th layer (Sun et al., 2016; Neyshabur et al., 2015; Xie et al., 2015). Although the sizes of the vectors corresponding to  $C_{\sigma}C_{w}$ can be exponentially large in depth, the norms of the vectors (and  $C_{\sigma}C_{w}$  itself) need not be (for example, by path-sparsity), and previous bounds also contain  $\prod_{\ell=1}^{H+1} C_{W^{(\ell)}}$  that can grow exponentially in depth (in addition to the explicit exponential dependence  $2^{H}$ ). By using a constant  $\alpha$ , our bounds can provide significantly improved asymptotic convergence rates when compared to the known rates (Role 2.2).

# 6 Theoretical Insight for Practical Usage

The previous two sections focused on theoretical bounds on generalization. However, in general, theoretical bounds can be too loose to be directly used in practice (even if they are tight in the sense of theoretical lower bounds. See Section 3.1). Accordingly, this section illustrates the use of generalization theory to provide theoretical insight to guide model class search in practice (Role 3).

In this section, we focus on multi-class classification problems with  $d_y$  classes, such as object classification with images. Accordingly, we analyze the expected risk with 0–1 loss as  $R[f] = \mathbb{E}_x[\mathbbm{1}\{f(x) = y(x)\}]$ , where  $f(x) = \operatorname{argmax}_{k \in \{1, \dots, d_y\}}(h_k^{(H+1)}(x))$  is the model prediction, and  $y(x) \in \{1, \dots, d_y\}$  is the true label of x (see Section 2.4.1 in Mohri et al. 2012 for an extension to stochastic labels).

#### 6.1 Theoretical Insight

An application of the result by Koltchinskii and Panchenko (2002) yields the following statement: given a fixed  $\rho > 0$ , for any  $\delta > 0$ ,

with probability at least  $1 - \delta$ , for all  $f \in F$ ,

$$R[f] \le \hat{R}_{m,\rho}[f] + \frac{2d_y^2}{\rho m} \mathfrak{R}'_m(F) + \sqrt{\frac{\ln \frac{1}{\delta}}{2m}},$$

where  $\mathfrak{R}'_m(F)$  is a model complexity defined as

$$\mathfrak{R}'_{m}(F) = \mathbb{E}_{S_{m},\xi} \left[ \sup_{k,h_{k}^{(H+1)}} \sum_{i=1}^{m} \xi_{i} h_{k}^{(H+1)}(x_{i}) \right].$$

Here,  $\xi_i$  are the Rademacher variables (i.e., independent uniform random variables in  $\{-1, +1\}$ ), and the supremum is taken over all  $k \in \{1, \ldots, d_y\}$  and all  $h_k^{(H+1)}$  allowed in F.

Previous theories (Sun et al., 2016; Neyshabur et al., 2015; Xie et al., 2015) characterize the generalization gap by the upper bounds on the model complexities via the norms of the weight matrices and exponential dependence on the depth  $2^H$ . A close look at the proofs reveals that the norms of weight matrices come from the Cauchy–Schwarz inequality, which can induce a very loose bound. Moreover, the exponential factor  $2^H$  comes from bounding the effect of nonlinearity at each layer, which can also induce a very loose bound, resulting in a gap between theory and practice. Avoiding the use of these possibly loose bounds seems to be challenging if we aim at Roles 1–3 at the same time.

By focusing on Role 3, this section proposes to solve this issue by directly approximating the model complexity  $\mathfrak{R}'_m(F)$ , instead of deriving a possibly too-loose bound on it (for worst possible measures). Here, the need for the approximation comes from the fact that  $\mathfrak{R}'_m(F)$  contains the expectation with unknown measure on dataset  $S_m$ . The approximation of  $\mathfrak{R}'_m(F)$  essentially reduces to the approximation of the expectation over the dataset  $S_m$ . In contrast to the worst-case bounds that motivated previous methods, the approximated  $\mathfrak{R}'_m(F)$  and whole generalization gap do not necessarily grow along with  $2^H$  and the norms of the weights, as desired.

## 6.2 Method

The theoretical insight in the previous section suggests the following family of methods: given any architecture and method, add a new regularization term for each mini-batch as

$$\text{loss} = \text{original loss} + \frac{\lambda}{\bar{m}} \hat{\mathbb{E}}_{S_m,\xi} \left[ \max_k \sum_{i=1}^m \xi_i h_k^{(H+1)}(x_i) \right],$$

where  $\hat{\mathbb{E}}_{S_m,\xi}$  is some approximation of  $\mathbb{E}_{S_m,\xi}$ ,  $\bar{m}$  is a mini-batch size and  $\lambda$  is a hyper-parameter. Importantly, the approximation  $\hat{\mathbb{E}}_{S_m,\xi}$  is only used for regularization purposes and hence needs not be precisely accurate (as long as it plays its role for regularization).

Table 1: Test error (%). A standard variant of LeNet (LeCun et al., 1998) and ResNeXt-29(16  $\times$  64d) (Xie et al., 2016) are used for MNIST and CIFAR-10, and compared with the addition of the studied regularizer.

Method	MNIST	CIFAR-10
Baseline	0.26	3.52
DARC1	0.20	3.43

For example, it can be approximated by populations generated by a generative neural network and/or an extra data augmentation process. For simplicity, we call this family of methods as Directly Approximately Regularizing Complexity (DARC).

In this paper, we evaluated only a very simple version of the proposed family of methods as a first step. That is, our experiments employed the following simple and easy-to-implement method, called DARC1:

loss = original loss + 
$$\frac{\lambda}{\bar{m}} \left( \max_{k} \sum_{i=1}^{\bar{m}} |h_k^{(H+1)}(x_i)| \right), (2)$$

where  $x_i$  is the *i*-th sample in the training mini-batch. The additional computational cost and programming effort due to this new regularization is almost negligible because  $h_k^{(H+1)}(x_i)$  is already used in computing the original loss. This simplest version was derived by approximating the expectation over  $S_m$  with the empirical distribution of the training data and the effect of the Rademacher variables via absolute values.

## 6.3 Experimental Results

We evaluated the proposed method (DARC1) by simply adding the new regularization term in equation (2) to existing standard codes for MNIST and CIFAR-10. For all the experiments, we fixed  $(\lambda/\bar{m})=$ 

Table 2: Test error ratio (DARC1/Base)

	MNIST (ND) mean stdv		MNIST		CIFAR-10	
	mean	$\operatorname{stdv}$	mean	$\operatorname{stdv}$	mean	$\operatorname{stdv}$
Ratio	0.89	0.61	0.95	0.67	0.97	0.79

Table 3: Values of 
$$\frac{1}{m} \left( \max_k \sum_{i=1}^m |h_k^{(H+1)}(x_i)| \right)$$

Method	MNIST (ND)		MNIST		CIFAR-10	
	mean	$\operatorname{stdv}$	mean	$\operatorname{stdv}$	mean	$\operatorname{stdv}$
Base	17.2	2.40	8.85	0.60	12.2	0.32
DARC1	1.30	0.07	1.35	0.02	0.96	0.01

0.001 with  $\bar{m}=64$ . We used a single model without ensemble methods. The experimental details are in Appendix E.1. The source code is available at: http://lis.csail.mit.edu/code/gdl.html

Table 1 shows the error rates comparable with previous results. To the best of our knowledge, the previous state-of-the-art classification error is 0.23% for MNIST with a single model (Sato et al., 2015) (and 0.21% with an ensemble by Wan et al. 2013) and 3.47% for CIFAR-10 (Graham, 2014). Our new and simple regularization method improved the base models significantly enough to outperform the previous state-of-the-art results.

To further investigate the improvement, we ran 10 random trials with computationally less expensive settings, to gather mean and standard deviation (stdv). For MNIST, we used fewer epochs with the same model. For CIFAR-10, we used a smaller model class (preactivation ResNet with only 18 layers). Table 2 summarizes the improvement ratio: the new model's error divided by the base model's error. We observed the improvements for all cases. The test errors (standard deviations) of the base models were 0.53 (0.029) for MNIST (ND), 0.28 (0.024) for MNIST, and 7.11 (0.17) for CIFAR-10 (all in %).

Table 3 summarizes the values of the regularization term  $\frac{1}{m}(\max_k \sum_{i=1}^m |h_k^{(H+1)}(x_i)|)$  for each obtained model. The models learned with the proposed method were significantly different from the base models in terms of this value. Interestingly, a comparison of the base cases for MNIST (ND) and MNIST shows that data augmentation by itself implicitly regularized what we explicitly regularized in the proposed method.

## 7 Conclusion

By embracing a logically expected difference between theory and practice, we separated several roles for generalization theory and made contributions in each area: an explanation for generalization in deep learning (via Role 1), an improvement of generalization bounds (Role 2), and a useful theoretical insight for regularization (Role 3).

Our observations in Section 5.3 suggest that if we can properly handle the dependence of deep-path activation vectors  $z_i$  in an analysis, we can obtain a tight guarantee. Our two-phase training procedure made the theoretical analysis easier by explicitly breaking the dependence. However, it will be interesting to see if future work can handle the dependence tightly in analysis without using the two-phase procedure.

Our theoretical insight in Section 6 led to a family of new generalization methods, DARC. Our empirical evaluation of its simplest version showed several promising results for further investigation of DARC.

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## Appendix A.

## A Additional notation

The usual empirical margin loss  $\hat{R}_{m,\rho}[f]$  is defined as  $\hat{R}_{m,\rho}[f] = \frac{1}{m} \sum_{i=1}^{m} \ell_{\text{margin},\rho}(f(x_i), y_i)$ , where  $\ell_{\text{margin},\rho}$  is defined as follows:

$$\ell_{\text{margin},\rho}(f(x),y) = \ell_{\text{margin},\rho}^{(2)}(\ell_{\text{margin},\rho}^{(1)}(f(x),y))$$

where

$$\ell_{\mathrm{margin},\rho}^{(1)}(f(x),y) = h_y^{(H+1)}(x) - \max_{y \neq y'} h_{y'}^{(H+1)}(x) \in \mathbb{R},$$

and

$$\ell_{\mathrm{margin},\rho}^{(2)}(z) = \begin{cases} 0 & \text{if } \rho \le z\\ 1 - z/\rho & \text{if } 0 \le z \le \rho\\ 1 & \text{if } z \le 0. \end{cases}$$

We use the convention that the training model output  $\hat{Y}$  and the training target Y in matrix form are real m by  $d_y$  matrices, where  $d_y$  is the dimension of the targets. The training input matrix X is real m by  $d_x$  matrix, where  $d_x$  is the dimension of the input. For test dataset, we define  $\hat{Y}_{\text{test}}, Y_{\text{test}}$ , and  $X_{\text{test}}$  in the same way with the size of test dataset  $m_{\text{test}}$ .

For any matrix M, let Col(M) and Null(M) be the column space and null space of M.

## B Appendix for Section 3

## B.1 Theorem 8

Theorem 8 (traditional theory can fail to explain generalization of over-parameterized linear models) Consider a linear model with the training output matrix  $\hat{Y} = \Phi w$  where  $\Phi$  is a fixed input future of X. Let  $\hat{Y}_{\text{test}} = \Phi_{\text{test}} w$  where  $\Phi_{\text{test}}$  is a fixed input future of  $X_{\text{test}}$ . Let  $M = [\Phi^{\top}, \Phi^{\top}_{\text{test}}]^{\top}$ . Let  $w^*$  be the ground-truth parameters. Then, if n > m and if  $\text{rank}(\Phi) \geq m$  and rank(M) < n.

- (i) For any target  $Y \in \mathbb{R}^{m \times d_y}$ , there exist parameters w such that  $\hat{Y} = Y$ , and
- (ii) For any  $\epsilon, \delta \geq 0$ , there exist parameters w and paris of training dataset  $(\Phi, Y)$  and test dataset  $(\Phi_{\text{test}}, Y_{\text{test}})$  such that
  - (a)  $\hat{Y} = Y + \epsilon A$  for some matrix A with  $||A||_2 \le 1$ , and
  - (b)  $\hat{Y}_{test} = Y_{test} + \epsilon B$  for some matrix B with  $||B||_2 \le 1$ , and
  - (c)  $||w||_2 \ge \delta$  and  $||w w^*||_2 \ge \delta$ .

**Proof** Since  $\operatorname{rank}(\Phi) \geq m$  and  $\Phi \in \mathbb{R}^{m \times n}$ , the set of its columns span  $\mathbb{R}^m$ , which proves statement (i). Let  $w^* = w_1^* + w_2^*$  where  $w_1^* \in \operatorname{Col}(M^T)$  and  $w_2^* \in \operatorname{Null}(M)$ . For statement (ii), set the parameter as  $w := w_1^* + \epsilon C_1 + \alpha C_2$  where  $\operatorname{Col}(C_1) \subseteq \operatorname{Col}(M^T)$ ,  $\operatorname{Col}(C_2) \subseteq \operatorname{Null}(M)$ ,  $\alpha \geq 0$  and  $C_2 = \frac{1}{\alpha}w_2^* + \bar{C}_2$ . Since  $\operatorname{rank}(M) < n$ ,  $\operatorname{Null}(M) \neq \{0\}$  and there exist non-zero  $\bar{C}_2$ . Then,

$$\hat{Y} = Y + \epsilon \Phi C_1$$
,

and

$$\hat{Y}_{\text{test}} = Y_{\text{test}} + \epsilon \Phi_{\text{test}} C_1.$$

By setting  $A = \Phi C_1$  and  $B = \Phi_{\text{test}} C_1$  with a proper normalization of  $C_1$  yields (a) and (b) in statement (ii) (note that  $C_1$  has an arbitrary freedom in the bound on its scale because its only condition is  $\text{Col}(C_1) \subseteq \text{Col}(M^T)$ ). At the same time with the same parameter, since  $(w_1^* + \epsilon C_1) \perp C_2$ ,

$$||w^*||_2^2 = ||w_1^* + \epsilon C_1||_2^2 + \alpha^2 ||C_2||_2,$$

and

$$||w - w^*||_2^2 = ||\epsilon C_1||_2^2 + \alpha^2 ||\bar{C}_2||_2^2$$

which grows unboundedly as  $\alpha \to \infty$  without changing A and B, proving (c) in statement (ii).

#### B.2 Corollary 9

Corollary 9 (traditional wisdom can fails to explain generalization of linear models) If  $n \leq m$  and if  $\operatorname{rank}(M) < n$ , statement (ii) in Theorem 8 holds.

**Proof** It follows the fact that the proof in Theorem 8 uses the assumption of n > m and  $\operatorname{rank}(\Phi) \geq m$  only for statement (i).

## B.3 Proof of Proposition 1

**Proof** Consider statement (i). Given such a F, consider any pair  $(\mathcal{A}, S_m)$  such that  $\mathcal{A}$  takes F and  $S_m$  as input and outputs  $f_{\epsilon}^*$ . Clearly, there are many such pairs  $(\mathcal{A}, S_m)$ . For example, fix  $S_m$  and then fix  $\mathcal{A}$  such that  $\mathcal{A}$  takes F and  $S_m$  as input and outputs  $f_{\epsilon}^*$  (which already proves the statement), or even  $f_{\epsilon}^* + \delta$  where  $\delta$  becomes zero by the right choice of hyper-parameters and of small variations of F (e.g., architecture search in deep learning) such that F still satisfy the condition in the statement. This proves statement (i).

Consider statement (ii). Given any dataset  $S'_m$ , consider a look-up algorithm  $\mathcal{A}'$  that always outputs  $f^*_{\epsilon}$  if  $S_m = S'_m$ , and outputs  $f_1$  otherwise such that  $f_1$  is arbitrarily non-robust and  $|\ell(f^*_{\epsilon}(x), y) - \ell(f_1(x), y)|$ 

is arbitrarily large (i.e., arbitrarily non-stable). This proves statement (ii).

Note that while A' in the above proof suffices to prove statement (ii), we can also generate other non-stable and non-robust algorithms by noticing the essence of this proof discussed in the main text right before Proposition 1 as a rephrase of the proposition.

## B.4 Additional Explanation for Role of Generalization Theory

Role 2.2 is important and has been a more focus in theoretical literature. However, practical deep learning approach may change a model class flexibly as m increases in the way that is not considered in theory. For example, theory with the focus on Role 2.2 would conclude that the generalization gap is upper bounded by  $O(\sqrt{n/m})$  with n being the number of parameters, and hence approaches to zero as m increases. However, n may increase as m increase in practice, while still producing a good generalization gap. Thus, we believe that Role 2.1 is also important as well as Role 2.2.

## C Appendix for Section 4

## C.1 Proof of Proposition 2

**Proof** Consider a single fixed  $f \in F_{\text{val}}$ . Since  $F_{\text{val}}$  is independent from the validation dataset,  $z_{f,1}, \ldots, z_{f,m_{\text{val}}}$  are independent zero-mean random variables, given a fixed  $f \in F_{\text{val}}$ . Thus, we can apply Bernstein inequality, yielding

$$\mathbb{P}\left(\frac{1}{m_{\text{val}}}\sum_{i=1}^{m_{\text{val}}} z_{f,i} > \epsilon\right) \le \exp\left(-\frac{\epsilon^2 m_{\text{val}}/2}{\gamma^2 + \epsilon C/3}\right).$$

By taking union bound over all elements in  $F_{\text{val}}$ ,

$$\mathbb{P}\left(\bigcup_{f \in F_{\text{val}}} \left\{ \frac{1}{m_{\text{val}}} \sum_{i=1}^{m_{\text{val}}} z_{f,i} > \epsilon \right\} \right)$$

$$\leq |F_{\text{val}}| \exp\left(-\frac{\epsilon^2 m_{\text{val}}/2}{\gamma^2 + \epsilon C/3}\right).$$

By setting  $\delta = |F_{\rm val}| \exp\left(-\frac{\epsilon^2 m_{\rm val}/2}{\gamma^2 + \epsilon C/3}\right)$  and solving for  $\epsilon$  (via quadratic formula),

$$\epsilon = \frac{2C\ln(\frac{|F_{\rm val}|}{\delta})}{6m_{\rm val}} \pm \frac{1}{2}\sqrt{\left(\frac{2C\ln(\frac{|F_{\rm val}|}{\delta})}{3m_{\rm val}}\right)^2 + \frac{8\gamma^2\ln(\frac{|F_{\rm val}|}{\delta})}{m_{\rm val}}}.$$

By noticing that the solution of  $\epsilon$  with the minus sign results in  $\epsilon < 0$ , which is invalid for Bernstein inequality,

we obtain the valid solution with the plus sign. Then, we have

$$\epsilon \le \frac{2C\ln(\frac{|F_{\text{val}}|}{\delta})}{3m_{\text{val}}} + \sqrt{\frac{2\gamma^2\ln(\frac{|F_{\text{val}}|}{\delta})}{m_{\text{val}}}},$$

where we used that  $\sqrt{a+b} \leq \sqrt{a} + \sqrt{b}$ . By tanking the negation of the statement, we obtain that for any  $\delta > 0$ , with probability at least  $1 - \delta$ , for all  $f \in F_{\text{val}}$ ,

$$\frac{1}{m_{\text{val}}} \sum_{i=1}^{m_{\text{val}}} z_{f,i} \le \frac{2C \ln(\frac{|F_{\text{val}}|}{\delta})}{3m_{\text{val}}} + \sqrt{\frac{2\gamma^2 \ln(\frac{|F_{\text{val}}|}{\delta})}{m}},$$

where 
$$\frac{1}{m_{\text{val}}} \sum_{i=1}^{m_{\text{val}}} z_{f,i} = R[f] - \hat{R}_{\text{val}}[f]$$
.

Note that  $F_{\text{val}}$  needs to be independent from validation dataset as defined in the main text, and otherwise the proof of Proposition of 2 breaks down.

## D Appendix for Section 5

We use the following lemma in the proof of Lemma 5.

Lemma 10 (Matrix Bernstein inequality: corollary to theorem 1.4 in Tropp 2012) Consider a finite sequence  $\{M_i\}$  of independent, random, self-adjoint matrices with dimension d. Assume that each random matrix satisfies that  $\mathbb{E}[M_i] = 0$  and  $\lambda_{\max}(M_i) \leq R$  almost surely. Let  $\gamma^2 = \|\sum_i \mathbb{E}[M_i^2]\|_2$ . Then, for any  $\delta > 0$ , with probability at least  $1 - \delta$ ,

$$\lambda_{\max}\left(\sum_{i} M_{i}\right) \leq \frac{2R}{3} \ln \frac{d}{\delta} + \sqrt{2\gamma^{2} \ln \frac{d}{\delta}}.$$

**Proof** Theorem 1.4 by Tropp (2012) states that for all  $t \ge 0$ ,

$$\mathbb{P}\left[\lambda_{\max}\left(\sum_i M_i\right) \geq t\right] \leq d \cdot \exp\left(\frac{-t^2/2}{\gamma^2 + Rt/3}\right).$$

Setting  $\delta = d \exp\left(-\frac{t^2/2}{\gamma^2 + Rt/3}\right)$  implies

$$-t^2 + \frac{2}{3}R(\ln d/\delta)t + 2\gamma^2 \ln d/\delta = 0.$$

Solving for t with the quadratic formula and bounding the solution with the subadditivity of square root on non-negative terms (i.e.,  $\sqrt{a+b} \leq \sqrt{a} + \sqrt{b}$  for all  $a, b \geq 0$ ),

$$t \le \frac{2}{3}R(\ln d/\delta) + 2\gamma^2 \ln d/\delta.$$

## D.1 Proof of Proposition 4

**Proof** From the definition of induced matrix norm and the Cauchy-Schwarz inequality,

$$R[f_{\mathcal{A}(S_{m})}] - \hat{R}_{m}[f_{\mathcal{A}(S_{m})}]$$

$$\leq \sum_{k=1}^{d_{y}} \|\bar{w}_{\mathcal{A}(S_{m}),k}\|_{2}^{2} \lambda_{\max} \left( \mathbb{E}[zz^{\top}] - \frac{1}{m} \sum_{i=1}^{m} z_{i} z_{i}^{\top} \right)$$

$$+ 2 \sum_{k=1}^{d_{y}} \|\bar{w}_{\mathcal{A}(S_{m}),k}\|_{1} \left\| \frac{1}{m} \sum_{i=1}^{m} y_{ik} z_{i} - \mathbb{E}[y_{k}z] \right\|_{\infty}$$

$$+ \mathbb{E}[y^{\top}y] - \frac{1}{m} \sum_{i=1}^{m} y_{i}^{\top}y_{i}.$$

Then, the statement directly follows the conditions on  $S_m$  and  $\mathcal{A}$ .

## D.2 From Deep Models to Sum of Paths Expression

Here, we present a simple example for equation (1) in the main text. For fully-connected feedforward networks with ReLU and max pooling, we can write  $\sum_{\text{path}}$  in equation (1) simply as

$$\begin{split} & \sum_{\text{path}} \bar{x}_{\text{path}} \bar{\sigma}_{\text{path}}(x, w_{\sigma}) \bar{w}_{\text{path}, k} \\ & = \sum_{k_0 = 0}^{d_x} \sum_{k_1 = 1}^{n_1} \cdots \sum_{k_H = 1}^{n_H} x_{k_0} \bar{\sigma}_{k_1, \dots, k_H}(x, w_{\sigma}) \bar{w}_{k_1, \dots, k_H, k} \end{split}$$

where  $\bar{w}_{k1,...,k_H,k} = \left(\prod_{l=1}^{H+1} w_{k_{l-1}k_l}^{(l)}\right)$  is the product of weights for each path  $(w_{k_{l-1}k_l}^{(l)})$  is the weight between  $k_{l-1}$ -th unit and  $k_l$ -th unit at the l-th layer), and  $n_1,\ldots,n_H$  are the number of hidden unites at each layer. Here,  $\bar{\sigma}_{k_1,\ldots,k_H}(x,w_\sigma)$  represent ReLU nonlinearity and/or max pooling for each path. That is,  $\bar{\sigma}_{k_1,\ldots,k_H}(x,w_\sigma)=1$  if all the units in the path is active and  $\bar{\sigma}_{k_1,\ldots,k_H}(x,w_\sigma)=0$  otherwise.

## D.3 SGD Chooses Direction only in terms of $\bar{w}$ but not in terms of $w_{\sigma}$

Each step of (stochastic) gradient decent is the steepest decent direction in terms of the first order term of Taylor expansion of the objective function L at a current parameter w:

$$L(w + d) = L(w) + \nabla_w L(w)^T d + ||d|| g(w, d),$$

where  $\lim_{d\to 0} g(w,d) = 0$ . For sufficiently small ||d||, the first order term  $\nabla_w L(w)^T d$  becomes dominant,

which is minimized with (stochastic) gradient decent direction  $d = -\nabla_w L(w)$ . Recall that

$$h_k^{(H+1)}(x) = [\bar{x} \circ \bar{\sigma}(x, w_\sigma)]^\top \bar{w}$$

Note that  $\sigma(x, w_{\sigma})$  is 0 or 1 for max-pooling and/or ReLU nonlinearity. Thus, the derivative of  $[\bar{x} \circ \bar{\sigma}(x, w_{\sigma})]$  with respect to w is zero everywhere (except at the measure zero set where the derivative does not exists). Thus, by the chain rule (and power rule),  $\nabla_w L(w)^T d$  only contain the contribution from derivative of  $h_k^{(H+1)}$  with respect to  $\bar{w}$ , but not with respect to  $w_{\sigma}$ .

## D.4 Simple Implementation of Two-Phase Training Procedure

Directly implementing equation (1) requires the summation over all paths, which can be computationally expensive. To avoid it, we implemented it by creating two deep neural networks, one of which defines  $\bar{w}$  paths hierarchically, and another of which defines  $w_{\sigma}$  paths hierarchically, resulting in the computational cost at most (approximately) twice as much as the original cost. We tied  $w_{\sigma}$  and  $\bar{w}$  in the two networks during standard phase, and untied them during freeze phase.

For example, for ReLU nonlinearity  $\sigma_{ReLU}$ , the computation of the network can be re-written as:

$$\begin{split} h^{(l)}(x) &= \sigma_{\mathrm{ReLu}}(W^{(l)}h^{(l-1)}(x)) \\ &= \dot{\sigma}_{\mathrm{ReLu}}(W^{(l)}h^{(l-1)}(x)) \circ W^{(l)}h^{(l-1)}(x) \\ &= \dot{\sigma}_{\mathrm{ReLu}}(W^{(l)}_{\sigma}h^{(l-1)}_{\sigma}(x)) \circ W^{l}h^{(l-1)}(x) \end{split}$$

where  $W_{\sigma}^{(l)}=W^{(l)},\ h_{\sigma}^{(l-1)}=\sigma_{\mathrm{ReLu}}(W_{\sigma}^{(l)}h_{\sigma}^{(l-1)}(x))$  and  $\dot{\sigma}_{\mathrm{ReLu}}(M)_{ij}=0$  if  $M_{ij}<0$  and  $\dot{\sigma}_{\mathrm{ReLu}}(M)_{ij}=1$  if  $M_{ij}\geq0$ . Note that if  $W_{\sigma}^{(l)}=W^{(l)},$  we have that  $h_{\sigma}^{(l-1)}=h^{(l)}.$  In the two-phase training procedure, we created two networks for  $W_{\sigma}^{(l)}h_{\sigma}^{(l-1)}(x)$  and  $W^{(l)}h^{(l-1)}(x)$  separately. We then set  $W_{\sigma}^{(l)}=W^{(l)}$  during standard phase, and frozen  $W_{\sigma}^{(l)}$  and only trained  $W^{(l)}$  during freeze phase. Our source code is available at: http://lis.csail.mit.edu/code/gdl.html

#### D.5 Experimental detail in Section 5.3.1

For all MNIST(ND), MNIST and CIFAR-10, we used the same settings as those for Tables 2 and 3 in Section 6. That is, for all experiments, we still used the same fixed value of  $(\lambda/\bar{m})=0.001$  with  $\bar{m}=64$ . Other experimental detail is also identical to what is described in Appendix E.1, except that we used 1000 epochs for each standard and freeze phase.

#### D.6 Proof of Lemma 5

**Proof** We use the fact that for data pints  $i \in \{\alpha m + 1, \ldots, m\}$  (its size is  $m_{\sigma} = (1 - \alpha)m$ ),  $z_{\alpha m+1}, \ldots, z_m$  are independent random variables, because  $w_{\sigma}$  is frozen and independent from  $x_{\alpha m+1}, \ldots, x_m$ .

<u>For  $\beta_1$ :</u> Matrix Bernstein inequality (Lemma 10) states that for any  $\delta > 0$ , with probability at least  $1 - \delta/3$ ,

$$\lambda_{\max} \left( \mathbb{E}[zz^{\top}] - \frac{1}{m_{\sigma}} \sum_{i=1}^{m_{\sigma}} z_{i} z_{i}^{\top} \right)$$

$$\leq \frac{2C_{zz}}{3m_{\sigma}} \ln \frac{3d_{z}}{\delta} + \sqrt{\frac{2\gamma_{zz}^{2}}{m_{\sigma}} \ln \frac{3d_{z}}{\delta}}.$$

Here, Matrix Bernstein inequality was applied as follows. Let  $M_i = (\frac{1}{m_{\sigma}}A^{(i)})$ . Then,  $\sum_{i=1}^{m_{\sigma}}M_i = \mathbb{E}[zz^{\top}] - \frac{1}{m_{\sigma}}\sum_{i=1}^{m_{\sigma}}z_iz_i^{\top}$ . We have that  $\mathbb{E}[M_i] = 0$  for all i. Also,  $\lambda_{\max}(M_i) \leq \frac{1}{m_{\sigma}}C_{zz}$  and  $\|\sum_i \mathbb{E}[M_i^2]\|_2 \leq \frac{1}{m_{\sigma}}\gamma_{zz}^2$ .

For  $\beta_2$ : We apply Bernstein inequality to each  $(k, k') \in \{1, \ldots, d_y\} \times \{1, \ldots, d_z\}$  and take union bound over  $d_y d_z$  events, obtaining that for any  $\delta > 0$ , with probability at least  $1 - \delta/3$ , for all  $k \in \{1, 2, \ldots, d_y\}$ ,

$$\left\| \frac{1}{m_{\sigma}} \sum_{i=1}^{m_{\sigma}} y_{ik} z_i - \mathbb{E}[y_k z] \right\|_{\infty}$$

$$\leq \frac{2C_{yz}}{3m_{\sigma}} \ln \frac{6d_y d_z}{\delta} + \sqrt{\frac{\gamma_{yz}^2}{m_{\sigma}} \ln \frac{6d_y d_z}{\delta}}$$

For  $\beta_3$ : From Bernstein inequality, with probability at least  $1 - \delta/3$ ,

$$\mathbb{E}[y^\top y] - \frac{1}{m_\sigma} \sum_{i=1}^{m_\sigma} y_i^\top y_i \le \frac{2C_y}{3m} \ln \frac{3}{\delta} + \sqrt{\frac{2\gamma_y^2}{m} \ln \frac{3}{\delta}}.$$

#### D.7 Proof of Theorem 6

**Proof** Notice that there is a dependence between  $z_i$  and  $\bar{w}$  in the form of  $||g(z_i)|| ||\bar{w}||$  for some g, because  $\bar{w}$  is learned with all training dataset (instead of  $(1-\alpha)m$  sized partial dataset). This dependence is taken care of by the condition of the algorithm  $\mathcal{A}$  in the statement for the bounded norms. Unlike model complexity approach, the norms need not be bounded given the event where a dataset is not concentrated. From Lemma 5, we know that a event where dataset is concentrated happens with probability at least  $1-\delta$ . Thus, the norms are required to be bounded only conditioned on such a event, to ensure that

the occurrence of such an event immediately implies our bound. Hence, the statement directly follows Proposition 4 and Lemma 5.

#### D.8 Proof of Theorem 7

**Proof** Recall the following fact: using the result by Koltchinskii and Panchenko (2002), we have that for any  $\delta > 0$ , with probability at least  $1 - \delta$ , the following holds for all  $f \in F$ :

$$R[f] \le \hat{R}_{m_{\sigma},\rho}[f] + \frac{2d_y^2}{\rho m_{\sigma}} \mathfrak{R}'_{m_{\sigma}}(F) + \sqrt{\frac{\ln \frac{1}{\delta}}{2m_{\sigma}}},$$

where  $\mathfrak{R}'_{m_{\sigma}}(F)$  is Rademacher complexity defined as

$$\mathfrak{R}'_{m_{\sigma}}(F) = \mathbb{E}_{S_{m_{\sigma}},\xi} \left[ \sup_{k,h_k^{(H+1)}} \sum_{i=1}^{m_{\sigma}} \xi_i h_k^{(H+1)}(x_i) \right].$$

Here,  $\xi_i$  is the Rademacher variable, and the supremum is taken over all  $k \in \{1, \ldots, d_y\}$  and all  $h_k^{(H+1)}$  allowed in F. Then, for our parameterized model classes with any frozen  $w_{\sigma}$ ,

$$\mathfrak{R}'_{m_{\sigma}}(F)$$

$$= \mathbb{E}_{S_{m_{\sigma}},\xi} \left[ \sup_{k,\bar{w}_{k}} \sum_{i=1}^{m_{\sigma}} \xi_{i} [\bar{x}_{i} \circ \bar{\sigma}(x_{i}, w_{\sigma})]^{\top} \bar{w}_{k} \right]$$

$$\leq \mathbb{E}_{S_{m_{\sigma}},\xi} \left[ \sup_{k,\bar{w}_{k}} \left\| \sum_{i=1}^{m_{\sigma}} \xi_{i} [\bar{x}_{i} \circ \bar{\sigma}(x_{i}, w_{\sigma})] \right\|_{2} \|\bar{w}_{k}\|_{2} \right]$$

$$\leq C_{w} \mathbb{E}_{S_{m_{\sigma}},\xi} \left[ \left\| \sum_{i=1}^{m_{\sigma}} \xi_{i} [\bar{x}_{i} \circ \bar{\sigma}(x_{i}, w_{\sigma})] \right\|_{2} \right].$$

Because square root is concave in its domain, by using Jensen's inequality and linearity of expectation,

$$\mathbb{E}_{Sm_{\sigma},\xi} \left[ \left\| \sum_{i=1}^{m_{\sigma}} \xi_{i} [\bar{x}_{i} \circ \bar{\sigma}(x_{i}, w_{\sigma})] \right\|_{2} \right]$$

$$\leq \left( \mathbb{E}_{Sm_{\sigma}} \sum_{i=1}^{m_{\sigma}} \sum_{j=1}^{m_{\sigma}} \mathbb{E}_{\xi} [\xi_{i} \xi_{j}] [\bar{x}_{i} \circ \bar{\sigma}(x_{i}, w_{\sigma})]^{\top} [\bar{x}_{j} \circ \bar{\sigma}(x_{j}, w_{\sigma})] \right)^{1/2}$$

$$= \left( \sum_{i=1}^{m_{\sigma}} \mathbb{E}_{Sm_{\sigma}} \left[ \left\| [\bar{x}_{i} \circ \bar{\sigma}(x_{i}, w_{\sigma})] \right\|_{2}^{2} \right] \right)^{1/2}$$

$$\leq C_{\sigma} \sqrt{m_{\sigma}}.$$

Putting together, we have that  $\mathfrak{R}'_m(F) \leq C_{\sigma} C_w \sqrt{m_{\sigma}}$ .

## E Appendix for Section 6

## E.1 Experimental detail in Section 6.3

For MNIST:

We used the following fixed architecture:

- Layer 1 Convolutional layer with 32 filters with filter size of 5 by 5, followed by max pooling of size of 2 by 2 and ReLU.
- Layer 2 Convolution layer with 32 filters with filter size of 5 by 5, followed by max pooling of size of 2 by 2 and ReLU.
- Layer 3 Fully connected layer with output 1024 units, followed by ReLU and Dropout with its probability being 0.5.
- Layer 4 Fully connected layer with output 10 units.

Layer 4 outputs  $h^{(H+1)}$  in our notation. For training purpose, we use softmax of  $h^{(H+1)}$ . Also,  $f(x) = \operatorname{argmax}(h^{(H+1)}(x))$  is the label prediction.

We fixed learning rate to be 0.01, momentum coefficient to be 0.5, and optimization algorithm to be (standard) stochastic gradient decent (SGD). We fixed data augmentation process as: random crop with size 24, random rotation up to  $\pm 15$  degree, and scaling of 15%. We used 3000 epochs for Table 1, and 1000 epochs for Tables 2 and 3.

## For CIFAR-10:

For data augmentation, we used random horizontal flip with probability 0.5 and random crop of size 32 with padding of size 4.

For Table 1, we used ResNeXt-29(16  $\times$  64d) (Xie et al., 2016). We set initial learning rate to be 0.05 and decreased to 0.005 at 150 epochs, and to 0.0005 at 250 epochs. We fixed momentum coefficient to be 0.9, weight decay coefficient to be  $5 \times 10^{-4}$ , and optimization algorithm to be stochastic gradient decent (SGD) with Nesterov momentum. We stopped training at 300 epochs.

For Tables 2 and 3, we used pre-activation ResNet with only 18 layers (pre-activation ResNet-18) (He et al., 2016). We fixed learning rate to be 0.001 and momentum coefficient to be 0.9, and optimization algorithm to be (standard) stochastic gradient decent (SGD). We used 1000 epochs.